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LMEDLINE coverage updated

SISSARVA enhanced with complete althor names

CHEMCATS accession numbers revised

CAYCAPLUS enhanced with trench and German abstracts

CAYCAPLUS enhanced with French and German abstracts

CAYCAPLUS patent coverage enhanced

USDATFULL/USPATT enhanced with Tre reclassification

USDATFULL/USPATT enhanced with Tre reclassification

USDATFULL/USPATT enhanced with Tre experimental property tags

FSTA enhanced with new thesaurus edition

CAYCAPLUS enhanced with additional kind codes for granted patents

CAYCAPLUS enhanced with CAS indexing in pre-1907 records

Full-text patent databases enhanced with prodefined patent family display formats from INPADOCDB

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CAS FEGISTRY enhanced with additional experimental spectral property data

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FORIS renamed to SOFIS

INNADOCDB enhanced with monthly SDI frequency

CAYCAPLUS enhanced with printed CA page images fron 1867-1938

CAPIUS coverage extended to include traditional medicine patents

EMMASE, FMBAL, and LEMBASE reloaded with enhancements Welcome to STN International NEWS 13 AUG 20 NEWS 14 AUG 27 NEWS 15 AUG 27 NEWS 16 AUG 28 NEWS 16 SEP 13 NEWS 19 SEP 13 NEWS 20 SEP 17 NEWS 21 SEP 17 NEMS 21 SEP 17 CAplus coverage extended to include the patents patents

NEMS EXPRESS 19 SEPTEMBER 2007: CURRENT MINDOWS VERSION IS V8.2, CURRENT MACINTOSH VERSION IS V6.0c(EMG) AND V6.0Jc(OP), AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.

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Enter NEWS followed by the item number or name to see news on that specific topic.

<12/04/2007> Erich Leese

10/513699



G1 C.N

G2 OH, C, H, O, Ak, MeO, EtO, n-Pro, i-PrO, n-BuO, i-BuO, s-BuO, t-BuO

Structure attributes must be viewed using STN Express query preparation.

=> 5 11 full FULL SEARCH INITIATED 17:39:44 FILE 'REGISTRY' FULL SCREEN SWARCH COMPLETED - 22712 TO ITERATE

100.0% PROCESSED 22712 ITERATIONS SEARCH TIME: 00.00.01 276 ANSWERS

276 SEA SSS FUL L1

s» file caplus COST IN U.S DOLLARS SINCE FILE TOTAL FULL ESTIMATED COST 172.10 172,31

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=> d ll L1 HAS NO ANSWERS

<12/04/2007> Erich Leese

They are available for your review at:

http://www.cas.org/infopolicy.html

=> s 12 full L3 23 L2

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I.3 ANSMER 1 OF 23
ACCESSION NUMBER:
DOCUMENT NUMBER:
1007:675422 CAPLUS
1007:675422 CAPLUS
1007:675423 CAPL

DOCUMENT TYPE:

English FAMILY ACC. NUM. COUNT: PATENT INFORMATION,

> PATENT NO. KIND APPLICATION NO. DATE DATE US 2007142394 WO 2007075688

KG, PRIORITY APPLN. OTHER SOURCE(S): GI MARPAT 147:95554 Disclosed are compds. of the formula or a pharmaceutically acceptable salt thereof, and compns. and methods of treating obesity, metabolic syndrome and a cognition deficit disorder, alone or in combination with other agents. Compds. of formula I wherein a is n, 1 and 2; b is 0, 1, 2, 1 and 4; U and M are Cli or one of U and W is Cli and the other N, when M! is CR and (un)substituted alkyl, M2 is N; n is 1 and 2; p is 0, 1 and 2; X is a bond, alkylene, alkenylene, CO, O, CR2O, etc., Y is CR2, CCR2O, CO, CN-OR and derivs., S, SP, and SO2; when M! is N, M2 is N; n is 2; p is 1 and 2; X is bond, alkylene, alkenylene, CO, NHCO, CCC, SC and SO2; Y is CR2, (CR2)2, CO, c-NOR and derivs., S, SD and SO2; when M! is N, M2 is N; n is 2; p is 1 and 2; X is bond, alkylene, alkenylene, CO, NHCO, CCC, SC and SO2; Y is CR2, (CR2)2, CO, S-SO and SO2; when M! is N, M2 is CH; n is 1 and 2; p is 0, 1, and 2; X is bond, alkylene, alkenylene, CO, NHCO, COC, SC and SO2; Y is O, CR3, (CR2)2, CO, C-NOR and derivs., S, SO, and SO2; Z is bond, (un)substituted alkyl, (un)substituted (heterolaryl(alkyl), etc., each R3 is independently H, halo, (halo)alkyl, CR3, CCF3, NO2; COSH and derivs., NR2 and derivs., etc., R5 is H, halo, (halo)alkyl, (DR, alkow) and CR1, each R8 is independently H, alkyl, oli, alkowy, halo, CP3, CCF3, NO2; COSH and derivs., NR2 and derivs., etc., R5 is H, halo, haloalkyl, (un)substituted cycloalkyl, (un)substituted (heterolaryl, and acyl; and their pharmaceutically acceptable salts thereof, are claimed. Example compound II was prepared by a multistep procedure (procedure given). All the invention compds were evaluated for their histamine H3 antagonistic activity (data given).

94270-93-1P

RL FAC (Pharmacological activity); SPN (Synthetic preparation); USES (USE)

(USES)

(drug candidate; preparation of substituted aniline derivs. as histamine H3 antagonists u

<12/04/2007>

Erich Leese

OTHER SOURCE(s): MARPAT 146,109356

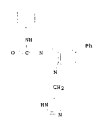
AB The invention provides methods for treating symucleinopathies, e.g. Parkinson's disease, diffuse Lewy body disease, and multiple system atrophy, comprising administering a symucleinopathic subject a farnesyl transferase inhibitor.

IT 195982-03-7

RE: PRG (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study), USES (Uses) (farnesyl transferase inhibitors for treatment of symucleinopathies)

RN 195982-03-7 CAPLUS

CN 4H-14-Benzodiasepine-4-carboxamade, 1,2,3,5-tetrahydro-1-(1H-imidazol-5-ylmethyl)-N-1-naphthalenyl-7-phenyl-, hydrochloride (1:1) (CA INDEX NAME)



● HC1

LJ ANSMER J OF 23 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMBER:
DOCUMENT NUMBER.
116: Le4488
Preparation of piperazinomethyl substituted
quinazolines useful in cancer treatment
Mallams, Alan K., Dammahapatra, Bimalendur, Neustadt,
Bernard R., Demma, Mark; Vaccaro, Henry A.
Schering Corporation, USA
Schering Corporation, USA
POCUMEN; Typs.
DOCUMEN; Typs.
LANGUAGE;
FAMILY ACC, NUM, COUNT:
English
FAMILY ACC, NUM, COUNT:
English
FAMILY ACC, NUM, COUNT:
English

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO KIND DATE APPLICATION NO.

PATENT NO. KIND DATE APPLICATION NO. DATE

MO 2007011623 A1 2007015 WC 2005-US27114 20060713

M: AE. AG. AL. AM. AT. AU, AZ. BA, BB, BG, BR, BW, BY, BZ, CA, CH,
CN, CO, CK, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
GE, GH, GM, HN, HR, HH, 10, I, I, IN, IS, JP, KE, KG, KM, KN, KP,

Erich Leese

DATE

PAGE 1-A

L3 ANSWEK 2 OF 23 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2007:133786 CAPLUS DOCUMENT NUMBER: 146:309356

2007;133786 CAPLUS
146:309356
Methods using farnesyl transferase inhibitors for the treatment of synucleinopathies
Lansbury, Peter T., Liu, Zhihua
The Brigham and Nomen's Hospital, Inc., USA
Aust. Pat. Appl., 520pp.
CODEN: AUXXCM
Patent
English

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. DATE AU 2006230674 PRIORITY APPLN. INFO.: AU 2006-230674 AU 2006-230674 Al 20061116 20061018

<12/04/2007>

Erich Leese

10/513699

GR, HU, IE, TR, BF, BJ, TG, BW, GH, AM, AZ, BY,

The title compds. [[m = 0-2, X = 0KS, N(R6)2; K1, R2 = H, alkyl; R3 - (un)substituted alkyl, cycloalkyl, aryl, etc.; R1 - alkyl; R4 - alkyl, cycloalkyl, aryl, etc.; R3 - alkyl; R4 - alkyl; cycloalkyl, aryl, etc.; R5, K6 = H, alkyl, cycloalkyl, etc.], useful for treating cellular proliferative diseases, disorders associated with activity of mutants of p53, or in caussing apoptosis of cancer cells, were prepared E.g., a multi-sitep synthesis of II, starting from Et 2-aminobenzoate and chloroacetonitrile, was given. Compound II showed RC50 of 1.1 µM (MB468) when tested in proliferation assay measuring the growth suppression effects of small mols. in cells with mutant p51 vs. p53 null background. The present invention also provides compns. comprising the compds. 1. 922153-20-8F 922156-06-7F 922159-12-4F RE: PAC (Pharmacological activity); SFN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (USES)

es)
(preparation of paperazinomethyl substituted quinazolines as antitumor

agents)
922151-20-6 CAPLUS
L-Valine, N-[2-[[4-[(1-naphthalenylamino)carbonyl]-1-piperazinyl]methyl]-4quinazolinyl]-. methyl ester (CA INDEX NAME)

<12/04/2007>

Erich Leese

Absolute stereochemistry.

Ο,

9:2156-06-7 CAPLUS
1-Piperazinecarboxamide, 4-[[4-[[flS]-1-(aminocarbonyl)-2-methylpropyl]amino]-2-quinasolinyl]methyl]-N-1-naphthalenyl-NAME] (CA INDEX

Absolute stereochemistry

0 NH2



922159-12-4 CAPLUS
1-Piperazinecarboxamide, 4-[(4-[[3-(dimethylamino)propyl]amino]-2quinazolinyl|methyl]-N-1-naphthalenyl- (CA INDEX NAME)

<12/04/2007.

Erich Leese

10/513699

US 2005-695306P WO 2005-CH5017 CASREACT 145:62919, MARPAT 145:62919 P 20050630 W 20051222 OTHER SOURCE(S)

R3 0

(Uses)
(preparation of alkoxybenzenecarboxamides as poly(ADP-ribose)polymerase (PARP) inhibitors (or the treatment of cancel)
87(83)-20-8 CAPUUS
1-Piperazinecarboxamide, 4-[5-[[2-(aminocarbonyl)-4-fluorophenoxy]methyl]-2 fluorobenzoyl]-N-1-naphthalenyl- (CA INDEX NAME)

10/513699

REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE PORMAT

L3 ANSWER 4 OF 23 ACCESSION NUMBER: 2006:605658 CAPLUS

DOCUMENT NUMBER: 116:62919

TITLE: 2006:605658 CAPLUS

145:62919

Preparation of 2-alkoxybenzenecarboxamides as poly(ADP-ribose)polymerase (PARP) inhibitors for the treatment of cancer

Javaid, Muhammad Hashim, Smith, Graeme Cameron Murray, Martin, Nall Morrison Barr, Gomez, Sylvie: Loh., Vincent Junior Ming Lai: Cockcroft, Xiao-Ling Fan, Menear. Keith Allan

PATENT ASSIGNEE(S): SOURCE: U.S. Pat. Appl. 101. 41 pp.

COURN: USXXCO
PALENT

DOCUMENT TYPE: LANGUAGE: English

FINILY ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

			DATE				ATE
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US 2006	135770	A1	20060622	US 2005-	315528	2	0051222
WO 2006	067472	A1	20060629	WO 2005-	GB5017	2	0051222
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			CZ, DE, DK,				
			MU, ID, IL,				
			LS. LT. LU.				
			NO, NZ, OM,				
			SY, TJ, TM,				
	VN, YU,			,,,			,
RW:			CY, CZ, DE,	DK. RE. RS.	PT. FR.	GR. GR.	HU. TE
			LV, MC, NL,				
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	KG, KZ.			30, 34, 12,	0G, ZM,	ZH, MM,	AL, B1
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				118 2004 -	638912P	P 2	0041223

<12/04/2007>

Erich Leese

10/513699

PAGE 1-A

PAGE 2-A

L3 ANSHER 5 OF 23 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMBER: 2006:117041 CAPLUS
DOCUMENT NUMBER: 144:212800
TITLE: Preparation of piperidine and

ALTIVES
144:212800
Preparation of piperidine and piperazine derivatives as histamine H3 receptor liquads for treatment of depression
Folmer, James, Hunt, Simon Praser, Hamley, Peter, WesolowSki, Steven
Astrazeneca AB, Swed.
PCT Int. Appl., 67 pp.
CODEN- PIXXD2
Patent
English
1

INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE .

LANGUAGE: FAMILY ACC, NUM, COUNT: PATENT INFORMATION:

<12/04/2007>

DATE APPLICATION NO PATENT NO. DATE WO 2006014135 2006014135 A1 20060209 WO 2005-5E1168 20050727 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN 2005-80026233 IN 2007-DN231 US 2007-572966 NO 2007-1140 SE 2004-1970 WO 2005-SE1188 OTHER SOURCE(S): WO 2005-SE1188 CASREACT 144:212800, MARPAT 144:212800

R N 1 N N AF Q Ar

The title piperidine and piperazine derivs, with general formula of 1 and II (wherein R = alky), 0 = N(CH2CH2) 2CH-, N(CH2CH2) 2CH-, UNICH2CH2) 2CH-O-, N(CH2CH2) 2CH-O-, N(CH2CH2) 2CH-O-, N(CH2CH2) 2CH-O-, NCH2CH2) 2CH-O-, N(CH2CH2) 2CH-O-, NCH2CH2) 2CH-O-, NCH2CH2 2CH-

are useful in the apy, in particular in the treatment of depression (no data)
875546-37-5P 875546-61-5P
Rt: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological atudy); PREP (Preparation); USRS (Uses)
(drug candidate; preparation of piperidine and piperazine derivs, as histamine H3 receptor ligands for treatment of depression)
875546-37-5 CAPLUS
1-Piperazinecarboxamide, N-(5-amino-1-naphthalenyl)-4-methyl- (CA INDEX

<12/04/2007>

Erich Leese

10/513699

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

IN 200527272 Al 20051208 US 2005 84739 20050318
PRIORITY APPLM. 1NFO. US 2004-55071P P 20040318
OTHER SOURCE(S): MARPAT 143:339666
AM Methols are provided of troating synucleinopathies, such as Parkinson's disease, diffuse Levy body disease and multiple system atrophy. Comprising admin.storing tr a synucleinopathic subject a farnesyl transferase inhibitor compound
IT 19582-03-7
KL. PAC (Pharmacological admin.storing)

195982-03-7

KL. PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USKS (Uses) (farnesy) transferase inhibitors for treatment of synucleinopathies) 195982-03-7 (AP-LUS 4H 1,4-Benzodrarepine-4-carboxamide, 1,2,3,5-tetrahydro-1-(IH-imidazol-5-ylmethyl)-N-1-naphthalenyl-7-phenyl-, hydrochloride (1:1) (CA INDEX NAME)

[],[]0- - C - N CHo

• HCl

10/513699

NAME)

875546-61-5 CAPLUS
1-Piperazinecarboxamide, 4-methyl-N-(5,6,7,8-tetrahydro-1-naphthalenyl)-(CA INDEX NAME)

REFERENCE COUNT: THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 6 OF 23 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2005:1049851 CAPLUS

2005:1043861 (APLOS 143:339666 Methods using farnesyl transferase inhibitors for the treatment of symucleinopathies Lansbury Peter T.; Liu, Zhihua The Brigham and Women's Hospital, Inc., USA PCT Int. Appl., 205 pp. CODEN: PIXKO2 DOCUMENT NUMBER: TITLE;

INVENTOR(S): PATENT ASSIGNEE(S); SOURCE:

DOCUMENT TYPE: LANGUAGE: Patent English

<12/04/2007> Erich Leese

10/513699

L3 ANSWER 7 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
111.256034
Preparation of phthalazinones as PARP inhibitors
Martin, Niall Morrison Barr, Smith, Graeme Cameron
Murray, Jackson, Stephen Philip, Loh, Vincent M., Jr.
Cockcroft, Xiao-Ling Fan; Matthews, Ian Timothy
Williams, Menear, Keith Allan, Kerrigan, Frank,
Abboorth, Alan
Kudos Pharmaceuticals Lamited, UK, Maybridge Limited
PCT Int. Appl., 102 pp.
COOMENT TYPE:
DOCUMENT TYPE:
Patent

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

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CA	2517	629			A1		2004	0923									
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IN	2005	DNO3	895		A		2007	0427		IN 2	005-1	8 E MC	95		2	0050	BJI
ZA	2005	0070	97		А		2006	0628		ZA 2	005-	70 97				0050	
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										MO 3	004-	GBIO	59		A 2	0040	312

OIHER SOURCE(S): MARPAT 141;296034

NH 0 · N N RΊ p11 p12 r

The title compds [I; A and B together represent (un)substituted fused aromatic ring; λ = NRx or CRxRy; if X = NRx then n = 1 or 2 and if X = CRxRy then n = 1, Rx = H. (un)substituted C1-20 alkyl, C5-20 aryl, C3-20 heterocyclyl, amado. thioamido, ester, acyl, and sulfonyl groups; Ry = H, OH. NH2; or Rx and Ry may together form a spiro(c3-7)cycloalkyl or heterocyclyl group; R11 and R12 are both H, or when X = CRxRy, R11, R12, RX and Rv. together with the carbon atoms to which they are attached, may form (un)substituted fused aromatic ring; R1 = N, halo), were prepared Thus, reacting 3-(4-oxo-3-4-dihydrophthalazni-1-ylmethyl)benoic acid (preparation given) with tert-Bu 1-piperazinecarboxylate afforded 77% II which had IC50 of < 0.02 yM against PARP. All compds. I tested had a IC50 of < 0.1 µM in the PARP assay. The pharmaceutical composition comprising the compound I is claimed.

µm in the PARP assay. The pharmaceutical composition comprising the I is Claimed. 763113-44-6P RE: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use), BIOL (Biological study); PREP (Preparation); USES (USES)

(preparation of phthalazinones as PARP inhibitors) 763113-44-6 CALLUS

763113-44-6 CALLUS
1-Piperacinecarboxamide, 4-[3-[(3,4-dihydro-1-oxo-1-phthaiazinyl)methyl]benzoyl]-N-1-naphthalenyl- (CA INDEX NAME)

'nн o-- c

REFERENCE COUNT: THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSHER 8 OF 23 CAPLUS COPYRIGHT 2007 ACS ON SYN ACCESSION NUMBER: 2004:612492 CAPLUS DOCUMENT NUMBER 141:156959

<12/04/200?> Erich Leese

10/513699

727724 - 96 - 1P RE: PAC (Pharmacological activity), SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(Uses)

(preparation of \$\beta\$-lactam compds. as tryptase inhibitors)

727'24-96-1 CAPLUS

2-Azetidinecarboxylic acid, 1-[[4-[(1-naphthalenylamino)carbonyl]-1-piperazinyl]carbonyl]-4-oxo-3-(4-piperidinylmethyl)-, (25.3R)- (CA INDEX NAME)

Absolute stereochemistry.



L3 ANSWER 9 OF ACCESSION NUMBER: DOCUMENT NUMBER: TITLE.

INVENTOR (S)

PATENT ASSIGNUE(S) :

DOCUMENT TYPE FAMILY ACC NUM, COUNT PATENT INFORMATION:

EP 1499607

<12/04/2007>

M3 2073048154

M AE. AG. AL.

CO. CR. CU.

MH. HU.

LE. LU.

PL. A. G.

RW: GH. CW.

KW: GH. CW.

FI. FR. GW.

CF. CG. CI.

CA 2473892

RP 1499607

Erich Leese

10/513699

Preparation of β-lactam compounds as inhibitors of tryptame Bisacchi, Gregory S.; Suttor, James C.; Slusarchyk, Milliam A.; Treuner, Uwe; Zhao, Guohua TITLE:

INVENTOR (S):

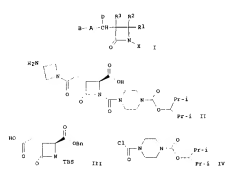
PATENT ASSIGNEE(s): SOURCE:

USA
U.S. Pat. Appl. Publ., 109 pp.
CODEN: USXXCO
Patent
English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

US 2003-728276 US 2002-434060P DATE US 2004147502
PRIORITY APPLN. INFO.;
OTHER SOURCE(S);
GI

MARPAT 141:156959



Beta lactam compds., such as 1 [R1 = H, carboxy, alkoxycarbonyl, alkenylaryl, CO-heterocyclyl, etc.; R2, R3 = H, alkyl; D = H, ORa; Ra = H, alkyl; D = GO-heterocyclyl, cycloheterocyclyl-Co, substituted amido, cycloalkyl, aryl, heteroaryl, cycloheteroalkyl; B = amino, aminoalkyl, aminocycloalkyl, cycloheteroalkyl, aryl, heteroaryl, alkylamino, carboxamidol, are prepared Thus, II was prepared Via a multistep synthetic sequence starting from [1-(diphenylmethyl)-3-azetidinyl)-carbamic acid-1;-1-dimethylethyl ester, III, and piperazinyl derivative IV. These compds. are useful as inhibitors of tryptase, thrombin, trypsin, Factor Xa, Factor VIIa, and urokinase-type glasminogen activator and may be employed in preventing and/or treating asthma and allergic rhinitis.

<12/04/2007> Erich Leese

10/513699

MX 2004PA06599
ZA 2004005348
US 2005043535
PRIORITY APPLN. INFO.: OTHER SOURCE(S): MARPAT 139:36450

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yllureido]ethyl]piperidine-4-carboxylic acid N-(naphthalen-1-yllamide Ri: PAC (Pharmacological activity); SFN [Synthetic preparation]; TNU (Therape-utic use); BIOL (Biological study); PREF (Preparation); USES

(Uses)
(urotensin antagonist; preparation of ureidoquinolines and analogs as urotensin 11 receptor antagonists for treatment of vasoconstriction, proliferation, and other disorders)
540769-67-3 CAPLUS
4-Pipertialnecarboxamide, 1-{2-{f[{2-methyl-4-quinolinyl}amino}carboxnyl}aminolethyl]-N-1-naphthalenyl- (CA INDEX NAME)

N-- CH2 CH2 NH C-- D 0- C

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT REFERENCE COUNT:

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ACCESSION NUMBER:

DOCUMENT NUMBER:

119-6892

Preparation of disaccycloalkane substituted piperatines as inhibitors and/or destabilizing androgen receptor ligans for the treatment of tumor illnesses. e.g. prostate cancer

Cleve, Arwed, Huwe, Christoph, Schulze, Volker; Morack, Helmut; Zopf, Dieter; Hoffmann, Jens, Reichel, Audreas

PATENT ASSIONEE(S): Schering Aktiengesellschaft, Germany

POCUMENT TYPE:

LANGUAGE: Patent

German

FAMILY ACC. NIM, COUNT: German

DOCUMENT TYPE-LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PAT	PATENT NO.					KIND DATE				APPL	ICAT:	ON	NO.		DATE		
															-		
WO	2003	04391	8.3		AI		2003	0530	,	WO 2	002-	EPI2	182		2	0021	031
	W:	AE.	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	ВB,	BG,	BR.	BY,	BZ,	CA,	CH,	CN,
		co,	CR,	cu,	CZ,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,
		HR,	HU,	ID.	IL,	IN.	IS,	JP,	KE,	KG,	KP.	KR,	KZ,	LC,	LK,	LR,	LS,
		LT,	LU,	LV.	MA,	MO,	MG.	MK,	MN,	MW,	MX,	MZ.	NO,	NZ,	OM,	PH,	PL,
		PT,	RO,	RU,	SD,	SE.	EG,	SI,	SK,	SL,	TJ.	TM,	TN,	TR,	TT,	TZ.	UA,
		UG,	UZ,	vc,	VN.	YU,	ZA,	ZM,	ZW								
	RW:	GH,	GM,	KE,	LS,	MW,	MZ.	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,
		KG,	rz,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE.	DK.	EE.	ES,
		FI,	FR,	GH,	GR,	1E.	IT,	LU,	MC.	NL,	PT,	SE,	SK,	TR,	BF,	ВJ,	CF,
		CG,	CI,	CM,	GA,	GN,	GQ,	G₩,	ML,	MR,	NE,	SN,	TD,	TG			
DE	1015	9035			7.1		2003	0612		DF 2	C01-	1015	9035		2	0011	123

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(CH2)4 ċ==0

PAGE 2-A

PAGE 1-A

534608-89-4 CAPLUS
1-Piperasinecarboxamide, 4-[5-[1-[4-cyano-3-(trifluoromethyl)phenyl]-2.5-dinydro-4-methyl-2.5-dioxo-1H-pyrrol-3-yl]pentyl]-N-1-naphthalenyl- (CA INDEX NAME)

10/513699

DE 10238742	A1	20040304	DE	2002-10238742		20020819
AU 2002360932	A1	20030610	ΑU	2002-360932		20021031
US 2004009969	A1	20040115	us	2002-301871		20021122
US 6861432	B2	20050301				
PRIORITY APPLN, INFO.:			DΕ	2001-10159035	A.	20011123
			DE	2002-10238742	A	20020819
			บร	2002-383785P	P	20020530
			US	2002-406650P	P	20020829
			WO	2002-EP12182	w	20021031
OTHER SOURCE(S):	MARPAT	139:6892				

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFELINE FRINT *

AB Title compds. I [A = CH3CO, CH3CONH, CN, etc., B = H, halo, CF3, etc., T = C or N with provisos; U = O, S; O = C(CH3)2, =CCH3)2, RI, R2 = H, CH3; i, j = 1-2, i+j = 2 or 3] their pharmaceutically acceptable salts and formulations were prepared For example, N-alkylation 1,2-dimethylethyl piperazin-1-carboxylate with iodopyrrol II, e.g., prepared from di-Me acctylene dicarboxylate with iodopyrrol II, e.g., prepared from di-Me acctylene dicarboxylate in 4-steps, provided claimed piperazine III. In inhibition of LNCAP cell proliferation, 18-examples of compds. I exhibited ICSO values ranging from 0.2-3 d x 10-7 M. Compds. I are claimed useful for the treatment of prostate cancer and benign prostatic hyperplasia.

IT 534609-10-19, 4 (-4.1-1.4-cyano-3-(trifuoromethyl))phenyl]-2,5-dihydro-4-methyl-2,5-dihydro-4-methyl-2,5-dioxo-1H-pyrrol-3-yllpepralin-1-carboxamide 534609-94-669, 4-[6.1-[4-Cyano-3-(trifluoromethyl)]phenyl]-3,5-dihydro-4-methyl-2,5-dioxo-1H-pyrrol-3-yllpepralin-1-vallpepralin-1-carboxamide 634609-94-69, 4-[6.1-[4-Cyano-3-(trifluoromethyl)]phenyl]-3,5-dihydro-4-methyl-2,5-dioxo-1H-pyrrol-3-yllpepralin-1-yllpperazin-1-carboxamide RD, PRAC (Pharmacological activity), PSPN (Synthetic preparation); TNO (Therapeutic usc), BGL (Biological study); PREP (Preparation); USES (BGL) DIA (BGL) (

<12/04/2007> Erich Leese

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FAGE 3-A (CH₂) 5

PAGE 2-A

DJ80UJ-96-6 CAPLUS
1-Piperazinecarboxamıde, 4-{6-{1-{4-cyano-3-(trifluoromethyl)phenyl}-2,5-dibydo-4-methyl-2,5-dioxo-1H-pyrrol-3-yl}hexyl]-N-1-naphthalenyl-INDEX NAME)

Erich Leese <12/04/2007> <12/04/2007> Erich Leese

PAGE 1-A

PAGE 2-A

REFERENCE COUNT: THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Į.

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ACCESSION NUMMER:
DOCUMENT NUMBER:
117:26:378

Novel potent antagonists of human neuropeptide Y YS receptor. Part 1: 2-oxobenzothiszolin-1-acetic acid derivatives

AUTHOK(SI:
Tabuchi, Selichiro; (tani, Hiromichi, Sakata, Yoshihiko; Cohashi, Hilzoko; Satoh, Yoshihari
FUSHINGE:
Bioorganic & Medicinal Chemistry Research Laboratories, Csaka, Yodoghwa-ku, 512-8514, Japan

SOURCE:
Bioorganic & Medicinal Chemistry Letters (2002), 112(8), 1171-1175
COERN: BMCLES; ISSN: 0960-894X

PUBLISHER:
Journal English

PUBLISHER: UDCUMENT TYPE: LANGUAGE: OTHER SOURCE(S):

English CASREACT 137:262978

<12/04/2007> Erich Leese

10/513699

Fujisawa Pharmaceutical Co., Ltd., Japan Jpn. Kokai Tokkyo Koho. 88 pp. CODEN: JKXXAF PATENT ASSIGNEE(S):

DOCUMENT TYPE:

Patent Japanese

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. PATENT NO. KIND DATE DATE JP 2001139574
PRIORITY APPLN. INFO.:
OTHER SOURCE(S):
GI A 20010522 JP 2000-296175 AU 1999-3093 20000928 MARPAT 134:366868

The title compds 1 [F1 \times H, halo: W \times S, O; A \times (CH2)n, etc.; n = 1 \times 6, Z \times (un)substituted N-containing hatercoyclic ring) are prepared 1-([5-Ch10c-2-oxobersothazolin-3-yllacevylpjpertdine-4-carboxylic acid 4-benoylamilide showed IC100 of 10-7 M in a neuropeptide Y5 receptor biodien each

4-benroylamilide showed fC100 of 10-7 M in a neuropeptide Y5 receptor binding 485ay.

J40179 71-4P 140178-83-8P
RL BAC (Biological actualty or effector, except adverse), BSU (Biological study, unclassified); SFM (Synthetic preparation); TBU (Therapeutic use); BIOL (Biological study); PREP (Breparation); USES (Uses)

JOURNAL OF THE PROPERTY OF THE P

s o o ci

34n178-81-8 CA:LUS 4-Piperidimecarboxamide, 1-[(5-chloro-2-oxo-3(2H)-benzothiazolyl)acetyl]-N-(5 hydroxy-1-napathalenyl)- (9CI) (CA IMDEX NAME)

10/513699

Novel neuropeptide NPY-YS antagonist FR7)966 I was discovered by screening of our inhouse chemical library. The analogs, e.g. II, were prepared by application of parallel synthesis techniques. Some of the resulting 2-oxobenzothiazolin-3-acetic acid derivs, exhibited nanomolar binding affinity for human NeY-YS receptors.

140178-71-8

RE: PRC (Pharmacological activity); SPM (Synthetic preparation); BIOL (Biological study), PREP (Preparation)
(preparation of 2-oxobenzothiazolin-3-acetic acid derivs, as potent antagonists of human neuropeptide Y YS receptor)

140178-71-4 CAPLUS
4-Piperidinecarboxamide, 1-[(S-chloro-2-oxo-3(2H)-benzothiazoly1)acety1]-N-1-naphthalenyl- (9CI) (CA INDEX NAME)

IT

REFERENCE COUNT: THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Li ANSHER 12 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2001;172159 CAPLUS
DOCUMENT MUMBER: 134;166868
TITLE: Preparation of benzothiazolines as neuropeptide Y receptor antagonists
SAUCY (Soshiya; Itani, Hiromachi, Tabuchi, Seiichiro, Sakata, Yoshihko; Ohashi, Hiroko

<12/04/2007>

10/513699

L3 ANSWER 13 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
2001:12274 CAPLUS
104:86272
Preparation of pyrimidine derivatives as Src-family
protein tyrosine kinase inhibitor compounds
llunt, Julianne A.; Mills, S.inder G.; Sinclair, Peter
J.; Zaller, Dennia M.
PATENT ASSIGNEE(S);
SOURCE:
DOCUMENT TYPE:

ACCOUNTEMENT TYPE:

COEN. PIXKD2

PATENT ASSIGNEE 20

COEN. PIXKD2

PATENT ASSIGNEE 30

COEN. PIXKD2

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: Patent English

Erich Leese

OTHER SOURCE(S); MARPAT 134:86272

<12/04/2007> Erich Leese <12/04/2007>

<12/04/2007>

10/513699

317365-53-0 CAPLUS
1-Piperazinecarboxylic acid, 2-[(1R)-1-[[4-(1H-benzimidazol-1-yl)-2-pyrimidinyl]amino]ethyl]-4-[(1-naphthalenylamino]carbonyl]-, phenylmethyl ester, (2S)-rel- (CA INDEX NAME)

Relative stereochemistry.

317365-56-3 CAPLUS
1-Piperazinecarboxylic acid, 2-[1-[[4-(1R-benzimidazoi-1-yl)-5-bromo-2-pyrimidinyl]amino]ethyl]-4-[[1-naphthalenylamino]carbonyl]-, phenylmethyl ester (CA INDEX NAME)

0 Ph CH2 O C N NH CH 11 N C- - 0 NH fi

317365-62-1 CAPLUS
J-Piperacinecarboxylic acid, 2-[1-[[2-(1H-benzimidazol-1-yl)-4pyrimidinyl]amino]ethyl]-4-[(1-naphthalenylamino)carbonyl)-. phenylmethyl
ester (CA IPDEX NAME)

10/513699

(Benzyloxycarbonyl)-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)ethylamino]-4-(indol-1-yl)pyrimidine 317365-76-7P, 2-[1-(1-(Benzyloxycarbonyl)-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)ethylamino]-4-[5-(3-ethylimidiacolidin-2-on-1-yl)benzimidazol-1-yl)pyrimidine 317365-80-1P, (s, s)-2-[1-(1-(Benzyloxycarbonyl)-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)ethylamino]-4-[5-(yyzidin-4-yl)benzimidazol-1-yl]pyrimidine 317365-85-8P, (s, s)-2-[1-(1-(Benzyloxycarbonyl)-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)ethylamino]-4-[5-(2-aminopyrimidin-4-yl)benzimidazol-1-yl]pyrimidine 317365-87-0P, (s, s)-2-[1-(1-(Benzyloxycarbonyl)-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)ethylamino]-4-[benzimidazol-1-yl]pyrimidine 317365-87-0P, (s, s)-2-[1-(1-(Benzyloxycarbonyl)-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)ethylamino]-4-[benzimidazol-1-yl]pyrimidine 32-(3-exatant) 52-(3-exatant) 52-(3-exat

317365-49-4 CAPLUS
1-Piperazimecarboxylic acid, 2-[(1R)-1-[[4-(1H-benzimidazol-1-yl)-2-pyrimidinyl]amino]ethyl]-4-[(1-naphthalenylamino]carbonyl]-, phenylmethyl ester, (2R)-rel- (CA INDEX NAME)

Relative stereochemistry.

<12/04/2007> Erich Leese

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317365-69-6 CAPLUS
1-Piperazinecarboxylic acid, 2-[1-[[4-(1H-indo]-1-y])-2pyrimidinyl]amino]ethyl]-4-[(1-naphthalenylamino)carbonyl]-, phenylmethyl
ester (CA INDEX NAME)

317365-76-7 CACLUS
1-PiperarineCartoxylic acid, 2-[1-[[4-[5-(3-ethyl-2-oxo-1-imidazolidinyl]-1-H-benzinidazol-1-yl]-2-pyrimidinyl]aminolethyl]-4-[(1-naphthalenylamino)carbonyl], phenylmethyl ester (CA INDEX NAME)

317365-80-3 CAPLUS
1-Piperazinecarboxylic acid, 4-[(1-maphthalenylamino)carbonyl]-2-[(18)-1-[[4-[5-(4-pyridinyl)-1H-bensimidazol-1-y]l-2-pyrimidinyl]amino]ethyl]-, phenylmathyl ester. (2S)- (CA INDEX NAME)

Erich Leese

Absolute stereochemistry



RN 317365-85-8 CAPLUS
CN 1-Piperaminecatroxylic acid, 2-{(15)-1-[|4-|5-(2-amino-4-pyrimidinyl)-lB-benimidAcol-1-y1]-2-pyrimidinyl]aminolethyl]-4-|(1-naphthalenylaminolcatbnyyl]-, phenylmethyl ester, (25)- (CA INDEX NAME)

Absolute stereochemistry.

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<12/04/2007> Erich Leese

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IT 317364-90-2P, 2- ([1-Methyl-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)methylaminol-4-(benzimidarol-1-yl)pyrimidine 317364-93-5P, (R.R.P.)-2-[1-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(benzimidarol-1-yl)pyrimidine 317364-96-8P, (R.R.P.)-2-[1-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(benzimidarol-1-yl)pyrimidine 317364-97-8P, (R.R.P.)-2-[1-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(benzimidarol-1-yl)pyrimidine 317365-96-3P, 2-(Benzimidarol-1-yl)pyrimidine 317364-97-8P, (R.R.P.)-2-(1-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(5-(3-thyl)pyrimidine 317365-11-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(5-(3-thyl)pyrimidine 317365-15-4P, (S.S)-2-(1-(Methyl-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(5-(3-thyl)pyrimidine 317365-16-5P, (S.S)-2-(1-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(S-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(S-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yl)tethylaminol-4-(N-naphth-1-ylcarbamoyl)piperazin-2-ylcarbamoylpiperazin-

10/513699

PAGE 2-A

N 17365-87-0 CAPLUS
N 1-Piperazinecarboxylic acid, 2-{(1s)-1-{[4-(1H-benzimidazol-1-yl)-2-pyrimidinyl]amino]ethyl]-4-{(1-naphthalenylamino)carbonyl}-, phenylmethyl ester, (2s)- (CA INDEX NAME)

Absolute stereochemistry.

IT 317365-10-9P, (R*,R*)-2-[1-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piperazin-2-yllethylaminol-4-[5-(3-ethylimidazolidin-2-on-1-yllbenzimidazol-1-yllpyrimidine
RL: PEP (Physical, engineering or chemical process); SFN (Synthetic preparation); TMU (Therapeutic use); BIOL (Biological study), PREP (Preparation) PROC (Process); USBS (Usea)
(preparation as inhibitor of Src-family protein tyrosine kinases and chromatog, resolution of)
RN 317365-10-9 CAPLUS
CN 1-Piperazinecarboxamide, 3-[(1R)-1-[[4-[5-(3-ethyl-2-oxo-1-imidazolidinyl)-1H-benzimidazol-1-yl]-2-pyrimidinyl]aminolethyl)-4-methyl-N-1-naphthalenyl-, (3R)-rel- (CA INDEX NAME)

Relative stereochemistry.

<12/04/2007> Erich Leese

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(2-aminopyridin-4-yl)benzimidazol-1-yl]pyrimidine 317365-95-0P,
2-[1-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piporazin-2-yl)ethylamino]-4-(5-(2-aminopyridin-4-yl)benzimidazol-1-yl]pyrimidine 317365-95-1P,
2-[1-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piporazin-2-yl)ethylamino]-4-[5-(pyridin-4-yl)benzimidazol-1-yl]pyrimidine 317365-97-2P,
2-[1-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piporazin-2-yl)ethylamino]-4-[5-(pyridin-4-yl)benzimidazol-1-yl]pyrimidine 317365-98-3P
317365-99-4P, 2-[1-(1-Methyl-4-(N-naphth-1-ylcarbamoyl)piporazin-2-yl]ethylamino]-4-[5-(2-aminopyrimidin-4-yl)benzimidazol-1-yl]-6-[2-methylamino]-4-[5-(2-aminopyrimidin-4-yl)benzimidazol-1-yl]-6-[2-methylamino]-4-[5-(2-aminopyrimidin-4-yl)benzimidazol-1-yl]-6-[2-methylamino]-4-[3-(2-aminopyrimidin-4-yl)benzimidazol-1-yl]-6-[2-(hydroxymethyl)bpenzimidin-4-yl)benzimidazol-1-yl]-6-[2-(hydroxymethyl)bpenzimidin-4-yl)benzimidin-4-yl)benzimidazol-1-yl]-6-[2-(hydroxymethyl)bpenzimidin-3-yl)benzimidin-3-yl)-6-[3-(hydroxymethyl)bpenzimidin-3-yl)-6-[3-(hydroxymethyl)bpenzimidin-3-yl)-3-(hydroxymethyl)bpenzimidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-yl)-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydroxymethyl)bproximidin-3-(hydrox

N NH CH2 N C O

RN 317364-93-5 CAPLUS
CN 1-Piperazinecarboxamide, 3-[(1R)-1-[(4-(1H-benzimidazol-1-yl)-2pyrimidinyl|amino|ethyl]-4-methyl-N-1-naphthalenyl-, (3R)-relNAME)

Relative stereochemistry.

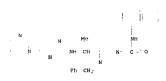
10/513699

317364-96-8 CAPLUS
1-0iperatinecarboxamide, 3-[(1R)-1-[(4-(1H-benzimidazoI-1-yI)-2pyrimidinyl]amino]ethyl]-4-methyl-N-1-naphthalenyl-, (3S)-rel- (CA INDEX NAME)

Relative stereochemistry,



317364-97-9 CAPLUS
1-Pipera:inecarboxamide, 3-{1-{{4-(IH-benzimidazo1-1-y1)-2-pyrimidinyl]amino|ethyl}-N-1-naphthalenyl-4-(plenylmethyl)- NAMB) (CA INDEX NAMB)



317365-06-3 CAPLUS
1-Piperarinecarboxamide, 3-{|-{{2-{1H-benzimidazol-1-y}};-4-}}
pyrimidinyl]amino|ethyl|-4-methyl-N-1-naphthalenyl- (CA INDEX NAME)

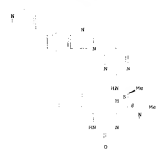
<12/04/2007>

Erich Leese

10/513699

317365-13-2 CAPLUS
1-Piperazinecarboxamide, 4-methyl-N-1-naphthalenyl-3-[(18)-1-[[4-[5-(4-pyridinyl)-1+benzimidxzol-1-yl]-2-pyrimidinyl]amino]ethyl]-, (38)- (CA JNDEX NAME)

Absolute stereochemistry. .



317)68-15-4 CAPLUS
1-Piperazinecarboxamide, 3-[(18)-1-[[4-{5-(2-amino-4-pyrimidiny1)-1H-benzimidazol-1-y|]-2-pyrimidiny1]amino]ethyl]-4-methyl-N-1-naphthalenyl-, (38)- (CA INDEX NAME)

Absolute stereochemistry

10/513699

317365-08-5 CAPLUS
1-Piperazinecarboxamide, 3-{1-[/4-(1H-indol-1-y1)-2-pyrimidinyl]aminolethyl]-4-methyl-N-1-naphthalenyl- (CA INDEX NAME)

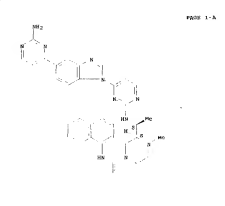
317365-11-0 CAPLUS
1-Piperarinecarboxamide, 3-{(18)-1-[4-{5-{3-ethyl-2-oxo-1-imidazolidinyl}}H-benzimidazol-1-yl}-2-pyrimidinyl]amino]ethyl] 4-methyl-N-1-naphthalenyl-, (3s)- (CA INDEX NAME)

Absolute stereochemistry.

<12/04/2007>

Erich Leese

10/513699



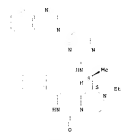
PAGE 2-A

317365-16-5 CAPLUS
1-Piperarinecarboxamide, 3-[(1S)-1-[[4-(1H-benzimidazol-1-yl)-2-pyrimidinyl]amino]ethyl]-4-methyl-N-1-naphthalenyl-, (3S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 317365-17-6 CAPLUS
CN 1-Piperarinecarboxamide, 3-[(1S)-1-[[4-(1H-benzimidazol-1-yl)-2-pyrimidinyl]amino]ethyl]-4-ethyl-N-1-naphthalenyl-, (3S)- (CA INDEX NAME)

Absolute stereochemistry.

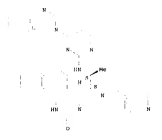


RN 317365-18-7 CAPLUS
CN 1-Piperazinecarboxamide, 3-[(1S)-1-|[4-(1H-benzimidazol-1-yl)-2pyrimidinyl]amino|ethyl]-4-hexyl-N-1-naphthalenyl-, (3S)- (CA INDEX NAME)

Absolute stereochemistry.

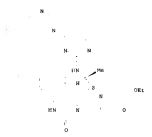
<12/04/2007> Erich Leese

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RN 317365-21-2 CAFLUS
CN 1-Papera.ineacetic acid, 2-[(18)-1-[[4-(]H-benzimidazol-1-yl)-2pyrimidinyl]am.nojechyl]-4-[(1-naphthalenylamino)cartonyl]-, ethyl ester,
(23)- (CA INDEX NAME)

Absolute stereochemistry



RN 317365-24-5 CAPLUS
CN 1-Piperarinecarboxamide, 4-acetyl-3-{(18)·1·{{4-(1H-benzimidazol-1-yl)·2-pyrimidinyl}amino|ethyl]-N-1-naphthalenyl, (38)· (CA INDEX NAME)

Erich Leese

Absolute stermochemistry

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10/513699

RN 317365-19-8 CAPLUS
CN 1-Piperazinecarboxamide, 3-[(18)-1-[(4-(1H-benzimidazo1-1-y1)-2pyrimidinyl|amino|ethyl]-4-(2-methylpropyl)-N-1-naphthalenyl-, (3S)- (CA
1NDEX NAME)

Absolute stereochemistry.

RN 317365-20-1 CAPLUS
CN 1-Piperacinecarboxamide, 3-[(15)-1-[[4-(1H-benzimidazol-1-yl)-2-pyrimidinyl]amino]ethyl]-N-1-naphthalenyl-4-(4-pyridinylmethyl)-, (38)-(CA INDEX MAME)

Absolute stereochemistry.

<12/04/2007> Erich Leese

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RN 317365-26-7 CAPLUS
CN 1-Piperazinecarboxamide, 3-{(1R)-1-[{4-|5-(3-ethyl-2-oxo-1-imidazolidinyl)1H-benzimidazol-1-yl]-2-pyrimidinyl]amino]ethyl]-4-methyl-N-1-naphthalenyl, (3R)- (CA INDEX NAME)

Absolute stereochemistry.

RN 317365-94-9 CAPLUS
CN 1-Piperazinecarboxamide, 3-{1-[{4-{5-{2-amino-4-pyzidinyi}-1H-benzimidazol-1-yl}-2-pyrimidinyl]amino|ethyl}-4-methyl-N-1-naphthalenyl- (CA INDEX NAME)

<12/04/2007> Erich Leese

317365-95-0 CAPLUS
1-Piperacinecarboxamide, 3-[1-[[4-[5-(2-amino-4-pyrimidinyl)-1H-benzimidazol-1-yl]-2-pyrimidinyl]amino]ethyl]-4-methyl-N-1-naphthalenyl-(CA INDEX NAME)



317365-96-1 CAPINS
1-Piperalinecarboxamide, 4-methyl-N-1-naphthalenyl-3-{1-{[4-{5-(4-pyridinyl)-14-benzimidazol-1-yl]-2-pyrimidinyl]amino]ethyl]- (CA INDEX NAME)

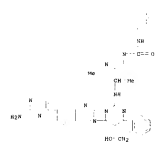


317365-97-2 CAPLUS
1-Piperaminecarboxamide, 4-methyl-N-1-naphthalenyl-3-[1-[[4-[5-(3-pyridixinyl)-lH-benzimidazol-1-yll-2-pyrimidinyl]amino]ethyl]- (CA INDEX NAMS)

<12/04/:007> Erich Leese

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31/36A-00-0 CAPLUS
1-P3pera:necarboxamide, 3-[1-[[4-[5-(2-amino-4-pyrimidinyl)-1H-benzimid:col-1-yl]-6-[2-(hydroxymethyl)ph-ny]}-2-pyrimidinyl]amino}ethyl]-4-methyl N-1-naphthalenyl- (CA INDEX NAME)



REFERENCE COUNT: THERE ARE 4 CITED REFFRENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSNER 14 OF 2) CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMBER: 2000:844922 CAPLUS
DOCUMENT NUMBER: 114:100734
TITLE: 114:100734
AUTHOR(S) Shuttlevorth, Stephen J; Nasturica, Daniel; Gervais,
Christian; Siddigui, M. Arshad, Rando, Robert F.; Lee,
Nola Biothem Sharma You Young Components

Nola BioChem Pharma Inc., Laval, QC, H7V 4A7, Can. Bioorganic & Medicinal Chemistry Letters (2000), 10(22), 2501-2504 COODN: BMCLER, TSSN: 0960-894X Elsevier Science Ltd. CORPORATE SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI Journal English CASREACT 134:100734



317365-98-3 CAPLUS
1-Piperazinecarboxamide, 3-[1-[[4-[5-[6-(dimethylamino)-3-pyridazinyl]-1H-benzimidazol-1-yl]-2-pyrimidinyl]amino]ethyl]-4-methyl-N-1-naphthalenyl-(CA INDEX NAME)

l17365-99-4 CAPLUS
1-Piperarinecarboxamide, 3-[1-[[4-[5-(2-amino-4-pyrimidiny1)-1H-benlimid20-1--yl]-6-(2-methylpheny1)-2-pyrimidinyl]amino]ethyl]-4-methyl-1-naphthalenyl- (CA INDEX NAME)

<12/04/2007> Erich Leese

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The synthesis of N-functionalized isatins, such as I (8 - CH (NelCCCGH4-3-OME, R1 - MC, R3 - H, R - CH3CCNUZ, CH3CCGH4-4-C1, R1 - R3 + H) suing parallel, solution synthesis is described. Punctionalized polymers were employed as stoichimmetric and catalytic reagents as well as purification media. The prepared isatins showed inhibition against a panel of serine proteases, i.e. human chymotrypsin, human ileukocyte elastase, and human plasmin.

119492-24-59 319492-26-79
RL: BAC (Biological activity or effector, except adverse), BSU (Biological study, unclassified), SPN (Synthetic preparation), BIOL (Biological study); PREP (Preparation)
(synthesis of isatin based serine protease inhibitors using polymer bound reagents)
119492-24-5 CAPLUS
1-Piperazinecarboxamide, 4-[[2,3-dihydro-2,3-dioxo-5-(trifluoromethoxy)-1H-indol-1-yl]acetyl]-N-1-naphthalenyl- (9CI) (CA INDEX NAME)

1T

l19492-26-7 CAPLUS
l-Piperazinecarboxemide, 4-{(5-chloro-2,3-dihydro-2,3-dioxo-1H-indol-1-yl)acetyl}-N-I-naphthalenyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT: THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 15 OF 2) CAPLUS COPYRIGHT 2007 ACS on STN
ACCRSSION NUMBER: 1999:690954 CAPLUS
DOCUMENT NUMBER: 1311:07106
Use of vitamin PP compounds as cytoprotective agents in chemotherapy
Biedermann, Elfi, Hasmann, Hax. Loser. Roland, Rattel,
Benno, Reiter, Fraedemain, Schein, Barbara,
Schemainda, Isabel, Seibel, Klaus, Vogt, Klaus,

Erich Leese

<12/04/2007> Erich Leese <12/04/2007>

Wosikowski, Katja Klinge Pharma GmbH, Germany PCT Int. Appl., 145 pp. CODEN: PIXXD2 PATENT ASSIGNME(S):

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

DE 1998-19818044 EF 1999-103814 WO 1999-FP2686 EP 2000-907642 WO 2000-EF1628 A 19980422 A 19990226 W 19990421 A3 20000228 W 20000228

OTHER SOURCE(S): MARPAT 131:307106

AB The invention relates to the use of vitamin PP compds. and/or compds. with anti-pellagra activity such as for example nicotinic acid (niacin), and nicotinamide (niacin-amide, vitamin PP, vitamin B3) for the reduction,

<12/04/2007> Erich Leese

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REFERENCE COUNT: THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 15 OF 23 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 1999:404932 CAPLUS

DOCUMENT NUMBER: TITLE:

131:58243

New piperazinyl-substituted pyridylalkame, -alkene, and -alkyne carboxamides, with antitumor and immunosuppressive activities

Biedermann, Elf; Hasmann, Max; Loser, Roland; Rattel,
Benno, Renter, Friedemann; Schein, Barbara; Seibel,
Klaus; Vogt, Klaus; Wosikowski, Katja
Klinge Pharma G.m.b.H., Germany
PCT Int. Appl., 224 pp.

CODEN: PIXXD2

Patent INVENTOR (S):

PATENT ASSIGNEE(S); SOURCE;

DOCUMENT TYPE:

Patent English

LANGUAGE: FAMILY ACC. NUM. COUNT PATENT INFORMATION:

PATENT N) KINU DATE APPLICATION NO. DATE

W) 9931063 A1 19990624 NO 1998-EP8268 19951216
W AL, AM, AT, AU, AZ, BA, BB, BG, PR, BY, CA, CH, CU, CZ, DE,
OK, FE, ES, FI, GB, GG, MG, MI, H, U, LD, LL, IN, IS, JF, KE,
KG, KF, KK, KZ, LC, LK, LR, LS, LT, LU, LU, MD, MG, MK, MN, MM,
MX, NG, NZ, PL, FT, RG, RU, SD, SE, SG, ST, KK, SL, TJ, TM, TM,
TT, UA, US, US, UZ, VN, 1U, ZW
RM: GH, GM, KE, LS, MM, SD, SZ, US, ZM, AT, BE, CH, CY, DE, DK, ES,
FI, FR, GB, GR, EE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CT,
CM, GA, GN, CM, ML, MM, E, SN, TD, TS
ZA 9811225 A1 19990701 DE 1997-1756236 19971217
ZA 9811225 A 19990608 ZA 1999-1756236 19991216
EP 1060163 A1 20001220 EP 1998-056275 19981216
EP 1060163 B1 20051012
R: AT, SE, CH, DE, DK, ES, FR, GB, CR, IT, LI, LU, NL, SE, MC, PT,
IE, F1
JP 2002508956 T 2002019 JP 2000-538990 19981276
ZA 20150478 T JP 2000-538990 AT 1998-965275 ES 1998-965275 US 2000-596001 DE 1997-19756236 WO 1998-EP8268 19981216 19981216 19981216 20000616 AT 306473 ES 2251794 US 6903118 PRIORITY APPLN. 1NFO.: A 19971217 W 19981216 OTHER SOURCE(S): MARPAT 131:58849

Erich Leese

10/513699

elimination or prevention of side-effects of different degrees as well as for neutralization of acute side-effects in immunosuppressive or cancerostatic chemotherapy or diagnosis, especially with substituted pyridine carboxamides, as well as combination medicaments with an amount of compds. With vitamin B3 and/or anti-pellagra activity and chemotherapeutic agents are especially considered in the mentioned chemotherapies and indications. Nicotinamide at 500 mg/kg twice daily protected mice treated i.p. with antitumor N-(4-(1-diphenylmethylpiperidin-4-yl)butyl]-3-(pyridin-3-yl)propionamide. There were no deaths in the nicotinamide-treated mice and the strong reduction of leukocytes was completely prevented. 227776-04-7 (Riological study): USES (Uses)
(Vitamin PP compds. as cytoprotective agents in chemotherapy) 227776-04-7 (ArLUS 1-P)personic Naturalization (N-1-naphthalenyl-4-(4-(1-oxo-3-(3-pyridinyl))rpopyl]amino|butyl]- (CA INDEX NAME)

PAGE I-A

PAGE 2-A

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$$\begin{array}{c|c} R^3 & \bigcap\limits_{\substack{i \\ A-C-N-D-S \cdots G \\ R^4 \\ R^1 \bigcap\limits_{\substack{i \\ 0 \\ 0}} N} \end{array}$$

The invention relates to new piperszinyI-substituted pyridylalkanoic,
-alkenoic, and alkynoic acid amides with a saturated or (poly)unsatd.
hydrocarbon residue in the carboxylic acid group, and analogs, i.e.,
haviug formula 1 [kl = H, OH, halo, cyano, COMH2, COCH, (heterolaryI,
alkoxy, amino, (heterolaryloxy, etc.; R2 = H, halo, cyano, alkyl, CF3, OH,
etc.; or R182 = (CH2)4, (CH:CH2)2, or CH2CCH20 or its (di)alkyl derivs, R3
= H, halo, alkyl, CF3, hydroxyalkyl, etc.; R4 = H, OH, alk(en/yn)yl,
cycloalkyl, alkoxy, aralkoxy, n = 0, 1; A = (unisubstituted alkylene or
hetero-isosteres, cycloalkylene, alkenylene, alkadienylene, or ethynylene,
D = (unisubstituted alkylene, alkenylene, alkynylene, or hetero-isosteres
of them, E = (unisubstituted (bis)(homo)piperazine bound at the N atoms; G
= variety of terminal chains]. Also disclosed are methods for the production
of the compds., medicaments containing them, and their production, as well as
their therapeutic use, especially as cytostatic agents and immuosuppressive
agents, for example, in the treatment or prevention of various types of
tumors, and control of immune reactions such as autoimmune diseases. For
example, 3-(3-pyridyl)acrylic acid was activated with oxalyl chloride and
condensed with 0-(3-(4-(diphenylmethyl)piperazin-I-yl)ropyl)hydroxylamine
to give title compound II. Several representative compds, inhibited various
human tumor cells in vitro at low concess e.g., with 1050 values of 0.1
nd to 10 µH, and also showed immunosuppressive activity against mouse
lymphocytes with 1050 values of 0.03-0,09 µM.

lymphocytes with IC50 values of 0.03-0,03 pM.
227776-04-7
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); USKS (Uses)
(target compound, preparation of piperazinyl-substituted pyridylalkanecarboxamides and analogs as cytostatics and immunosuppressants)
227776-04-7 CAPLUS

12:7/10-04-7 CAPLOS
1-Plperazinecarboxamide, N-1-naphthalenyl-4-{4-{{1-oxo-3-(3-pyridinyl)propyl]amino]butyl}- (CA INDEX NAME)

<12/04/2007> Erich Leese

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PAGE 1-A
 CHo
 CH2
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 NH
(CH<sub>2</sub>)<sub>4</sub>
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REFERENCE COUNT: THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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ANSWER 17 OF 23 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMMER: 1599;394051 CAPLUS
DOCUMENT NUMBER: 1199;394051 CAPLUS

INVENTOR(S): 11:44847
Preparation of heterocyclylbenzamidines as blood-coagulation factor %x inhibitors
Dorsch, Dieter, Jurasyyk, Horst, Wurziger, Hanns; Gante, Joachim, Mederski, Merner; Buchstaller, Hans-Peter, Annali, Soheila; Bernotat-Danielowski, Sabine, Melzer, Guido
Merck tatent G.m.D.H, Germany
Ger. Offen., 36 pp.
CODEN: GMXXBX
DOCUMENT TYPE
DOCUMENT TYPE
PATENT INFORMATION.

FAMILY ACC. NIM COUNT: 1
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<12/04/2007> Erich Leese

inhibitors)
227326-77-4 CAPLUS
1-Piperazinecartoxamide, 4-[(3-[4-(aminoiminomethyl)phenyl]-2-oxo-5-oxazolidinyl]methyl]-N-1-naphthalenyl-, monoacetate (9CI) (CA INDEX NAME)

CM 1

CRN 227326-76-3 CMF C26 H26 N6 03

PAGE 1-A

NHo ÇH2 c- o

PAGE 2 - A

CM 2

CRN 64-19-7 CMF C2 H4 O2

OTHER SOURCE(5):

R12I22CH2CH(OR3)CH223Z4R4 [R1 = (acyl- or hydroxy-substituted) C(:NH)NH2, 5-methyl-1,2,4-oxadiazol-3-yl, etc.: R3 = H, alkyl, CH2Ph, etc.: R4 = (cyclolalkyl, pienyl(alkyl), heterocyclyl(alkyl), etc.: Z1 = (un)substituted phenylene; Z2 = O or NR5, R5 = H, alkyl, CH2Ph, R3R5 = CO, Z3 = O, NR5, piperazine-1,4-diyl, etc.: Z4 = bond, CO, SO2, CO2, CONR5) were prepared as plood-coaylation factor Xa inhibitors (no data). Thus, 3-(4-(5-methyl-1,2,4-oxadiazol-3-yl)phenyll-2-oxooxazol(dine-5-ylmethyl methanesulfonate (preparation described) was aminated by Boc-piperazine and the deprotected product amidated by 2,4,6-trichlorobenzenesulfonyl chloride to give, after hydrogenation, title compound I.ROAC. 227326-77-49
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation) THU (Therapeutic use); BIOL (Biological study); PRE7 (Perparation); USPSS (USSS)
(preparation of heterocyclylbenzam:dines as blood-coayulation factor Xa

<12/04/2007> Erich Leese

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PAGE 1-A

PAGE 2-A

L3 ANSWER 18 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1999:233904 CAPLUS

DOCUMENT NUMBER:

130;282084
Benzamidine derivatives as factor Xa inhibitors
Dorsch. Dieter; Juraszyk, Horst; Murriger, Hanns;
Bernotat-Panielowski, Sabine; Melzer, Guido
Merck Patent G.m.b.H, Germany
PCT Int. Appl., 79 pp.
CODEN: PIXXD2
Patent
German
1 TITLE: INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PAT	ENT :	NO.			KIND)	DATE			APE	LI	CAT	ON	NO.				
WO	9916																9980	
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6.5	10,15																	
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		SI,	LT,	LV,	FI,	RO												
BR	9812 2001 2000	699			A		2000	0822		BR	19	98-	1269	9		1	9980	916
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<12/04/2007> Erich Leese

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CRN 64-19-7 CMF 62 H4 02 0

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REFERENCE COUNT. THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 19 OF 23 CAPLUS COPYRIGHT 2007 ACS On STN
SSION IMMNER; 1997.5/99715 CAPLUS
E: Imidazole-containing benzodiarcpines and analogs as inhibitors of farnesyl protein transferase
NIOF(S: Ding, Charles Z.: Hunt. John T.: Kim, Scong-hoon; ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: INVENTOR (S:

Erich Leese

10/513699

Title compds. I [X = bond, CO, (un)substituted CH2, CH2CH2, CH2CO, CH2CH2CO, CH:CHCO. NHCO; Y = (un)substituted CH3, SO2, CO, CO2, CONH; R = (un)substituted Ph; Rl = H, (un)substituted alkyl, oxaalkyl, thiaalkyl, alkenyl, cycloalkyl, aryl, aryloxy, heterocyclic, aralkenyl) are inhibitors of coagulation factor Xa and can be used for preventing or treating thromboembolic disorders (no data). Thus, 4-(5-methyl-1,2,4-oxadiazol-3-yl)benzoic acid was converted to the acid chloride, treated with N-tert.-butoxycarbonylpiperazine. and deblocked to give [4-(5-methyl-1,2,4-oxadiazol-3-yl)phenylpiperazin-1-ylmethanone which was treated with 6-chloro-2-naphthalenesulfonyl chloride and reduced to give the benzamidine II. 222541-47-7p
RL: STN (Synthetic preparation); USES (Uses)
(preparation of piperazinylbenzamidine derivs. as factor Xa inhibitors) 122541-47-7 CAPLUS
1-Piperazinecarboxamide, 4-(4-(aminoiminomethyl)benzoyll-N-1-naphthalenyl, monoacetate (SCI) (CA INDEX NAME)

CM 1

CRN 222543-46-6 CMF C23 H23 N5 O2

<12/04/2007> Erich Leese

10/513699

Mitt, Toomis; Bhide, Rajeev; Leftheris, Katerina Bristol-Myers Squibb Co., USA PCT Int. Appl., 425 pp. CODEN: PIXXD2 Patent PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: English

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					CH,	OE,	DK.	ES,	FR,	GB,	GR	, IT,	ы.	LU,	NL.	SE,	MC,	PT,	
			1E,	FI															
		1214				A		1999			CN	1997-	1925	35		1	9970		
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		1419				A		2003	0410		IL	1997- 1997- 1998- 1998- 2004-	1413	08			9970	224	
	11	1241 2225 4309	9/			Α		2003	0024		11	1391-	1241	9/			9970	224	
	KU	2225	405			62		2004	0310		KU	1998-	11//	98		1	9970	221	
	EE	1481				В1		2004	0.012		EB	1998-	1624	-		1	9970	224	
	BP	1481	9/5			V.T		2004	1201		EP	, IT.	1034	'		20,1	9970	224	
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		1211				B1		2006			PD.	1997- 1998-	1206	00			9970 9970	224	
		9701						1998			73	1007	1621			•	0070	224	
		4968				~		2002			TW	1997- 1997-	1021	2660		1	9970 9970	223 305	
		1215				В		1998				1998-					9980		
		9803				A		1998				1998-					9980		
	NO	3193	074			B1		2005			140	1990-	3092			-	,,,,	223	
		4552				8		1999			TT	1008-	120				9980	975	
		6495				B1		2006			PU.	1998- 1998- 1999-	1027	3.6		,	9980		
		6455				B1		2002			ite	1000	3742	10		1	9990		
		1347				A		2002											
		AHP A				^		2002	UJ 08		III O	1996-	1226	5 D		D 1	0010 9960	226	
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OTHER SOURCE(5): MARPAT 127:278213

<12/04/2007> Erich Leese AP The invention relates to a series of imidatole-substituted benzodiazepines and analogs that inhibit farnesyl-protein transferase (FPT) and ras protein farnesylation, thereby being useful as anti-cancer agents. The compds are also useful in the treatment of diseases, other than cancer, associated with signal transduction pathways operating through ras, and thos associated with proteins other than ras that are also post-translationally modified by FPT. The compds, may also act as inhibitors of other prenyl transferases, and thus be effective in the treatment of diseases associated with other prenyl modifications of proteins. Over 430 synthetic examples are given. For instance, 2,1,4,5-tectahydro-11-1,4-benzodiazepine was N-acylated by 1-naphthoic acid Ph ester in the presence of DMAP, and the product was reductively alkylated by 4-formylimidazole in the presence of NaHH(OAC)3 to give title compound f, isolated as the HCl salt. The example compds inhibited FPT with IC50 values between 0.1 nM and 100 µ.

IT 195982-01-TP xL: BMC (Biological activity or effector. except adverse), BSU (Biological study, unclassified), SPN (Synthetic preparation), TMU (Therapeutic une), BIO, (Biological study), PKEP (Preparation); USES (USES) (preparation of inhibitors

oitors
of (arnesyl protein transferase)
195982-03-7 CAPLUS
4H-1,4-Benzodiazepine-4-carboxamide, 1,2,3,5-tetrahydro-1-(1H-imidazol-5ylmethyl)-N-1-naphthaleny)-7-phenyl-, hydrochloride (1:1) (CA INDEX NAME)

<12/04/2007> Erich Leese

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OTHER SOUNCE(S): CASREACT 127:220673; MARPAT 127:220673

 $\left\|z^2\cdots\right\|_{\mathbb{R}^{d-1}\setminus\{0\}}$ R2

Title compds I [R1 = H. alky]; R2, R3 = H. alky], alkoxy, thioether, nitrile, CT3, F, Cl. Pr. I; or R2R1 form a 5- or 6-membered ring; XY = NCH2, CHCM2, CJCH, N. NCH2CM2, 21 = (CH2)H, (CR2)HCO, CO, CJCH2/HD, SO2, SO3/CHDH, OCH2)HO, OCC, NHCCJ2H, NICCH2)HO, CHCM1, CHCM2, CO, CHCM2/H, NHCCH2/SO2, NHSO2 (CH2)H, NHCCH2/SO2, NHSO2 (CH2)H, CH-CHCO, C.tplbond.cCO, (CM2)HH, H = 1-6, Ax1 = (unisubstituted Ph. Haphthyl, or pyridy); with provisos; are disclosed. The compds, are sirong and selective antagonists of 5 HTID receptors, and are useful for creatment of a variety of conditions, including depression, anxiety, schizophrenia, neurodep-merative disorders, and some cancers. Synthetic examples are quiven for 42 compds, and their fumarate salts. For instance, 4-methoxy-3-(4-methylpiperazin-1-yl)aniline underwent reaction with triphospune, and subsequent amidation with 4-phenethylpiperazine, to give 84% title compound II. In a test for inhibition of sumatripan-induced thymidine uptake by 6c glial cells transfected with the 5-HTIDB and 5-HTIDa receptor genes, I had 1C50 values in the range of 10-100 nM. In 5-HT receptor assays. II had Ki values of 2.1 nM and 1.9 nM for subtypes 1Da and 1DB, resp., vs. 3500 nM for subtype 1A. 194942-86-6P
RJ: BBC (Biological activity or effector, except adverse), BSU (Biological study): PREP (Preparation), USES (Uses) (preparation of piperazine derive, as 5-HTID antagonists)
194942-87-5 CAPLUS
14-Piperazinedicarboxtwide, N-[4-methoxy-3-(4-methyl-1-piperazinyl)phenyl]·N'·(5,6,7,8-tetrahydro-1-naphthalenyl) (9CI) (CA INDEX NAME)

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L3 ANSWER 20 OF 23 CAPLUS COPYRIGHT 2007 ACS ON STN

ACCESSION NUMBER: 1397:533632 CAPLUS

DOCUMENT NUMBER: 127:22073

TITLE: Comparison as method for preparing same, pharmaceutical compositions, and use thereof as drugs

HALARY, Serge, Jorand-Lebrun, Catherine, Pauwels, Peter; Chopin, Philippe, Marien, Marc

PATENT ASSIGNEE(S): Pierre Fabre Medicament, Fr., Halazy, Serge, Jorand-Lebrun, Catherine, Pauwels, Peter, Chopin, Philippe, Marien, Marc

SOURCE: Pierre Fabre Medicament, Fr., Halazy, Serge, Jorand-Lebrun, Catherine, Pauwels, Peter; Chopin, Philippe, Marien, Marc

POT Int. Appl., 131 pp.

COUNTY TYPE: Patent

LANGUNGE: Pierre

PAMILY ACC, NUM, COURT: 1

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PAT	ENT	NO.			KIN	D DATE		APP	LICATIO	N NO.		D	ATE		
	WO	9728	141			A1	1997	0807	WO	1997-FF	203		19	9702	203	
		₩:	ΑU,	BR,	CA,	CN,	JP, KR,	MX,	NZ, US							
		RW:	AT,	BE,	CH,	DE,	DK, ES,	FΙ,	FR, GB	, GR, 1	E, IT.	LU,	MC.	NL,	PT,	SE
	FR	2744	449			A1	1997	0808	FR	1996-12	73		15	99602	02	
	FR	2744	449			B1	1998	0424								
	CA	2245	718			A1	1997	0807	CA	1997-22	45718		15	9702	0.3	
	Aυ	9716	074			A	1997	0822	ΑU	1997-16	074		15	9702	03	
	EP	8805	12			A1	1998	1202	EP	1997-90	2427		19	9702	203	
		R:	AT,	RE,	CH,	DE.	DK, ES,	FR,	GB, GR	, IT, 1	ıI, LU,	NL,	SE,	MC,	PT,	
			IE,	FI												
	BR	9707	251			A	1999	0406	BR	1997-72	51		1	99102	03	
	CN	1214	047			A	1999	0414	CN	1997-19	3122		1:	99702	203	
	JΡ	2000	5057	95		T	2000	0516	JP	1997-52	7377		1	9702	03	
PRIO	RITY	APP	LN.	INFO					FR	1996-12	73		A 1:	99602	02	
									WO	1997-19	203	1	N 19	9702	203	

<12/04/2007> Erich Leese

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PAGE 1-A

PAGE 2-A

194942-88-6 CAPLUS
1,4-Piperazinedicarboxamide, N-[4-methoxy-3-(4-methyl-1-piperazinyl)phenyl]-N'-(5,6,7,8-tetrahydro-1-naphthalenyl)-,
(2E)-2-butenedioate (1:1) (9C1) (CA INDEX NAME)

CM 1

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PAGE 1-A 'nн Ċ-PAGE 2-A

CM 2 CRN 110-17-8 CMF C4 H4 O4

Double bond geometry as shown.

E со₂н H0 2 €

L3 ANSWER 21 OF 23 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1976;164871 CAPLUS
ORIGINAL REFERENCE NO: 48:267759,267798
EINVENTOKIG) Benzodiazepine derivatives
(NOVENTOKIG) Kindlarepine derivatives
(NOVENTOKIG) FAIGE, EVA, Szeberenyi, Szabolca; Szporny, Laszlo
PATENT ASSIGNAE(S): Richter, Gedeon, Vegyeszeti Gyar RT., (Nung.

<12/04/200% Erich Leese

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similar to that of diazepam against metrozole-induced convulsions, but with less sedative and muscle relaxant side effects and a much higher with less Secative
(DSO, 59010-23-0P

FRES PN (Synthetic preparation); PREP (Preparation)

(preparation of)

Spoilo-23-0 CAPIUS

4H-1,4-Benzodiazepine-4-carboxamide, 7-chloro-1,2,3,5-tetrahydro-1-methylN-1-naphthalenyl-2-oxo-5-phenyl- (SCI; (CA INDEX NAME)

 i_1,\dots,i_r NH Fh 0 . C . N | o

L3 ANSWER 22 OF 23 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMBER: 1961:87597 CAPLUS
DOCUMENT NUMBER: 155:87597
STRIGHTAL PROPERED NO. 55:187597-6
TITLE: 95:87597
TITLE: 155:18577-6
HOVED TITLE: 155:1857-6
HOVED T

ORIGINAL REFERENCE NO.: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): DOCUMENT TIPE: LANGUAGE: FAMILY ACC NUM, COUNT PATENT INFORMATION:

HATENT NO US 29/7362 KIND DATE US 29/7342 19610128 US 1957-703494 19571218

AM Acyl Werrys, of 1-(2-hydroxyechyl)-4-(N-phonylcarbamoyl)piperazine-HCl [I] were prepared by treating I with Rocol on (Roc)76 in the presence of a tertiary base or other acid acceptor. Thus, 10 g. I in 50 ml. pyridine was treated with 10 ml. Acyl, the mixture kopt ovenight at room temperature, distiller, and the residue dissolved in 20 ml. H20 and made basic with aqueous NH3. The mixture was extracted with Et20, the extract evaporated, and the less than a process of the control o APPLICATION NO. DATE due maice

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Ger. Offen., 48 pp. CODEN: GWXXEX Patent German 2 SCURCE: DCCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	KIND	DATE	APPLICATION NO.		DATE
DE 2523250	A1	19751218	DE 1975-2523250		19750526
DE 2523250	C2	19880107			
AU 7580951	A	19761111	AU 1975-80951		19750508
AU 502405	B2	19790726			
IL 47268	A	19810130	IL 1975-47268		19750512
CH 628036	A5	19820215	CH 1975-6729		19750522
FR 2272674	A1	19751226	FR 1975-16295		19750526
FR 2272674	B1	19790810			
SE 7506053	A	19751201	SE 1975-5053		19750527
SE 426242	В	19821220			
SE 426242	c	19830414			
BE 829595	A1	19750915	BE 1975-156798		19750528
DK 7502366	A	19751130	DK 1975-2366		19750528
DK 153479	В	19880718			
DK 153479	c	19881128			
NL 7506272	A	19751202	NL 1975-6272		19750528
JP 51001486	A	19760108	JP 1975-63985		19750528
DD 121516	A5	19760805	DD 1975-186307		1975052B
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PL 98943	Bl	19780531	PL 1975-193640		19750528
PL 100441	81	19781031	PL 1975-180778		19750528
CA 1063605	A1	19791002	CA 1975-227928		19750528
CS 195290	B2	19800131	CS 1975-3740		19750528
SII 942594	A3	19820707	SU 1975-2137707		19750528
SU 776559	A3	19801030	SU 1976-2343705		19760408
CS 195291	B2	19800131	CS 1977~178		19770111
JP 54055591	A	19790502	JP 1978-94608		19780713
JP 01022259	В	19890425			
SU 1318158	A3	19870615	SU 1978-2663501		19780918
RIORITY APPLN. INFO :			HU 1974-RI538	A	19740529
			CS 1975-3740		19750528

Benrodiazepines I (R = Cl, NO2, NH2, H; R1 = H, Me; R2 = alkoxy, amino, Cl. cycloalkyl. Me. CH2Cl, CH2NH2, CH2Ph, H, CH:CH2, C6H4Cl2 were prepared by treating 4-unsubstituted benrodiazepines with ClCOR2, isocyanates etc. I are tranquilizers. Thus I (R = Cl, R1 = Me, R2 = NH2) had a activity

<12/04/2007> Brich Leese

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228-98. Similarly, 1-(2-acetoxyethyl)-4(dimethylcarbamoyl)piperazine oxalate was prepared from AcCl and
1-(2-hydroxyethyl)-4-(dimethylcarbamoyl)piperazine (1V), m.
159-1609. IV was prepared by treating 10 g. II with 8-5 g. Me2NCCL
in CNCl3 at room temperature, and working up after 24 hrs. 1-(2-(NPherylcarbamoylcxy)ethyl)-4-(phrylcarbamoyl)-piperazine, prepared by
refluxing 5 g. free base of I with 2-7 ml. PhNco in 50 ml. CSH624 hrs., m.
180° (ECRH). Anticholinergic and antispasmodic properties were
shown by the salts of the new compds.
101578-29-4P, 1-riperazinecarboxamide, 4-(2-hydroxyethyl)-N-1naphthyl- 110441-89-9P, 1-Piperazinecarboxamide,
4-(2-hydroxyethyl)-N-1-naphthyl-, acetate, hydrochloride
RL: PREP (Preparation)
(preparation of)
101578-29-4 CAPLUS
1-Piperazinecarboxamide, 4-(2-hydroxyethyl)-N-1-naphthyl- (6CI) (CA INUEX
NAME)

но си2-си2

NH.

i

110441-89-9 CAPLUS
1-Piperazinecarboxamide, 4-(2-hydroxyethyl)-N-1-naphthyl-, acetate, hydroxhoride (fCI) (CA INDEX NAME)

Aco CH2 CH2 c ... o NH

● HC!

L3 ANSWER 24 OF 23 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION RUMBER.
DOCUMENT NUMBER.
DIGIONAL REPEWENCE NO : 46 6210a-1,6211a-d
TITLE
AUTHOR(S):
FOR FORTE SOURCE:
S 228 37 CODEN: BSCBAG: ISSN: 0037-9646 22.8 37
CODEN. BSCBAG: ISSN: 0037-9446

JOUTHAI
LANGUAGE

To diagram(s). see printed CA Issue.
AF The authors have prepared a large number of 4-substituted 1-methylpiperazines to be texted for filaricidal activity Products resulting from the reactions of 1-methylpiperazine (1) or its di-RCI salt with various reagents are subdivided into 4 categories: (1) reaction with amides or esters of halogenated aliphatic acids in alc. in the presence of NaHCO3 (2) reaction with acyl chlorides in various media in the presence of NaHCO3 (2) reaction with acyl chlorides in Tuesti). (3) reaction with various alkyl sulfouyl chlorides in CHCl3, (4) use of 1-methyl-4-piperazinecarbonyl chloride (III) in various media on a variety of amines (CA, 44, 3506g). To obtain crystallizable, nondeliquescent salts of these compds., various organic acids were used. Preparation of II: I 4 parts) in 100 parts PhMe at 0° is added with vigorous stirring to 20 parts (15% excess) COCl2 of the composition of the solvent of the solvent of the solvent who and dry ether (quant yields In certain cases, it saffices to remove them and trapped in NH3 solution Type preparation, category (1):

N,N-disthyl-1-methyl-1-1
piperazineacetamide. I 2HCl (346 g.) (2 mols), 504 g. (6 mols.) NaHCO3, and 50 g. (2 mols.) r CHCONNEZ are refluxed 10 h in 2.1 alc., the mixture cnilled, the mineral salts (altered oif, the solvent evaporated, dried by azoutropic distillation, the residue distilled under a high vacuum, 213 g.

c12/04/200%>

Erich Leese

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CON(CHIZCH2Ph)2. 215°/n 25. 70, 4, V. 186-8"; CONNCHIZCH2Ph, 170°/l. W8, 4, VI, 127°; CONMeRt, 192-4°/l5, 70, 4, thwarate, 130-1°; CONMeRt, 192-2, 70, 4, TV, 144°; CON CHIZ.CHIZ.CHIN, 138-40°/0,5, 90, 4, V1, 155°; CONICH.CHIC.CHIN, 128°/1,5, V0, 4, TV, 125°; CONICH.CHIC.CHIN, 128°/1,5, V0, 4, TV, 125°, CONICHHCLIC, N, 128°/1,5, V0, 4, TV, 125°, CONICHHCLIC, N, 128°/1,5, V0, 4, TV, 125°, CONICHC, NH)NH2, -, 50, 4, V1, decompose < 50°. VII was propared by saponification of the preceding ester in the presence of MeNH2, was was propared by action of 1-methyl-4-chloroacetopiperasine on EtzNH: IX was propared from the preceding amide.
6266-76-8P, 1-Piperazinecarboxamide, 4-methyl-N-1-naphthyl-85680-11-9P, 1-Piperazinecarboxamide, 4-methyl-N-1-naphthyl-hydrochloride
RP- PREP (Preparation)
(preptration of)
6266-76-8 CAPIUS ation of) CAPLUS

1-Piperazinecarboxamide, 4-methyl-N-1-naphthalenyl- (9CI) (CA INDEX NAME)

NH

856845-11-9 CAPLUS

1-Pip:razinecarhoxamide, 4-methyl-N-1-naphthyl-, hydrochloride (5CI) (CA INDEX NAME)

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mol.) of the base in 1.2 l. EtOH at 94° added to 232 g. (2 mols.) maleic acid in 1.2 l. EtOH at 94° with shaking: the dimaleate crystallizes immediately and quant. Type preparation, category (2): 1-Methyl-4-ethoxalyl piperazine. To 17.3 g. (0.1 mol.) I.2HCl and 12 g. KOH in 150 cc. HZO are added simultaneously with stirring, and at 0° 15 g. (0.11 mol.) ClCCOC2Et and 6 g. KOH in 10 cc. HZO stirring continued 1 h., the solution saturated with K2CO3, extracted with ether,

ether
extract dried, concentrated, distilled in a high vacuum, and the HCl salt
formed by
bubbling dry HCl into a C6H6-EtOH solution of the base; the product is easily
recrystd. from C6H6 or Me2CO containing 10% alc. Type preparation, category

N,N - Di-Et - 1 - Me - 4 - piperazinesulfonamide. Et2NSO2Cl (17.1 g. 0.1 mol.) in 50 cc. CRCl3 is added at 0* with stirring to 20 g. 10.2 mol.) I in 50 cc. CRCl3, the mixture refluxed 12 h., and the solvent removed at atmospheric pressure; the residue solidifies on cooling, and after

as atmospheric pressure; the residue solidifies on cooling, and after trituration with 10 cc. absolute alc., 100 cc. dry ether is added and the mixture filtered, giving 0.1 mol 1.1(Cl.) m. 130°. The filtrate is evaporated in vacuo, the oily product dissolved in 100 cc. ether, filtered with charcoal, and the ether solution added with agitation 0.1 mol. malelac acid in EtD-ECO. The product, an oil which crystallizes on scratching, is repptd. from a min. of absolute alc. with ether. Type preparation, category (4).

1-Methyl-4-(methylethylcarbamyllpiperazine. To 19.9 g. (0.1 mol.) II suspended in 100 cc. dry PhMe, at 0° is added with stirring 20 g. (0.3 mol.) EtMHMe in 100 cc. dry PhMe, at 0° is added with stirring 20 g. (0.3 mol.) StMHMe in 100 cc. dry PhMe, and the solution slowly heated and retluxed 1 h. The EtMHMe, HCl formed adheres to the walls of the flask. After cooling, 1 volume ether is added, the Et20-PhMe solution decanted, the solvent evaporated, and the residue distilled in vacuo. The fumarate is prepared by adding the base in ether to a suspension of 12 g, fumaric acid in 60

by adding the base in ether to a suspension of 12 g. fumaric acid in 60 cc. absolute alc., evaporating the ether, and the warming the alc. solution until the

until the salt dissolves; after crystallization 2 vols. ether is added and the salt repptd.

Table dissolves; after crystallization 2 vols. ether is added and the salt tid.

from a min. of 1so-PrON with ether. The following CH2.CH2.NNe.CH2.CN2.NNe. CH2.CN2.NNe. CH2.NNe. CH2.CN2.NNe. CH2.NNe. CH2.N

<12/04/2007>

Erich Leese

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Ŋ

● HCl

a) dhis DNIS IS NOT A RECOGNIZED COMMAND The previous command name entered was not recognized by the system. For a list of commands available to you in the current file, enter "NZLP COMMANDS" at an arrow prompt (+>).

=> d his

(FILE 'HOME' ENTERED AT 17:39:08 ON 17 NOV 2007)

FILE 'REGISTRY' ENFERED AT 17:39:16 ON 17 NOV 2007 STRUCTURE UPLOADED 276 S L1 FULL

FILE 'CAPLUS' ENTERED AT 17:39:50 ON 17 NOV 2037 23 S L2 FULL